## IN THE CLAIMS

- 1. (Currently Amended) A method of de novo designing molecules that bind to a receptor site on a protein comprising the steps of:
- (a) building a molecule in the receptor site comprising: adding successive random molecular fragments to an initial molecular fragment that is loaded into the receptor site, estimating the free energy of the molecule being grown after each addition of a molecular fragment, and orienting each successive molecular fragment as it is added to the receptor site such that the free energy estimate for the molecule may be higher than a lowest free energy estimate possible for the molecule;
- (b) repeating step (a) to generate a collection of molecules grown in the receptor site, and ranking the collection of molecules according to increasing free energy estimates to identify high-ranking molecules;
- (c) selecting one or more functional groups of a high-ranking molecule identified in step (b) as a single restart fragment and using the restart fragment to build a second-generation of molecules according to steps (a) and (b);
- (d) minimizing the energy of a protein/ligand complex comprising the receptor site and a second-generation molecule using an empirical force field;
- (e) quantitatively measuring the empirical interaction energy of the second-generation molecules, and ranking the <u>second-generation</u> molecules, wherein a <u>second-generation</u> molecule of low interaction strength is ranked higher than a <u>second-generation</u> molecule of more negative interaction energy is ranked higher than a <u>second-generation</u> molecule of less negative or positive interaction energy;
- (f) modifying high-ranking <u>second-generation</u> molecules from step (e) based on qualitative analysis of the <u>second-generation</u> molecules including determination of chemical viability, synthetic feasibility, solubility, and effect of the <u>second-generation</u> molecule on the structure of the protein, whereas such modification comprises: atomic and/or functional substitutions, initiating growth from a specific receptor site, inclusion of salt bridges or hydrogen bonds, and solubility-enhancing measures[.];
- (g) repeating steps (c) through (f) until a <u>second-generation</u> molecule is built which is identified as high-ranking in both steps (e) and (f)[.]; and

- (h) displaying at least one second-generation molecule built according to step (g).
- 2. (Original) The method of claim 1, wherein the receptor site is selected from the group consisting of: Src-homology-3 domain, Src-homology-2 domain, MDM2 protein, CD4 protein, and human carbonic anhydrase II protein.
- 3. (Original) The method of claim 1, wherein the empirical interaction energy comprises CHARMM interaction energy.
- 4. (Original) The method of claim 1, wherein the empirical force field comprises CHARMM.
- 5. (Cancelled).
- 6. (Cancelled).
- 7. (Cancelled).
- 8. (Cancelled).
- 9. (Cancelled).
- 10. (Cancelled).
- 11. (Cancelled).
- 12. (Cancelled).
- 13. (Cancelled).
- 14. (Cancelled).
- 15. (Cancelled).
- 16. (Cancelled).
- 17. (Cancelled).
- 18. (Cancelled).
- 19. (Cancelled).
- 20. (Cancelled).